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Studies in the Field of Organophosphorus Insecticides
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The present study was carried out in the laboratory of organophosphorus compounds under the direction of the Corresponding Member
of the Academy M.I.Kabachnik. The studies in the field of insecticides
were begun in 1952 and at first had a nonsystematic character, but
from 1953 and especially from 1954 numerous friganophosphorus compounds
were synthesized in our laboratory with the purpose of obtaining new
insecticides.

In 1953 there was established a constant contact with V.I.E.R. of V.A.S.Kh.N.I.L. where the tests of insecticidal properties of our compounds were made. This contact was expressed not only by the transfer of substances for tests but also, and this was very important, in a systematic discussion with the workers of V.I.Z.R. - D.M.Paikin, M.P.Shabanova and N.M.Gamper - of the results of the tests and means of preparing new active compounds. A similarly friendly contact we were able to establish with the representatives of the chemical industry. S.L.Varshavskii and S.V.Preobrashenskaya aided in many ways the successful completion of the present work. Part of the work was carried out in the laboratory of vinyl compounds of the Institute of Organic Chemistry of the Academy of Sciences (M.P.Shostakovskii, E.N.Prilezhawva) and in the laboratory for organic synthesis of V.N.I.I.V.M.P.T.Sh.P. (V.N.Odnoralova).

In order to have a possibility of selection among the uncountable number of emisting and possible organic substances of a definite and necessarily limited number of objects for the tests, it is necessary to be guided by some working hypothesis which permits one to carry out such a selection consciously and with aim.

In our studies we used the rather well accepted working hypothesis concerning the inhibition of cholinesterase as the cause of the toxic action of organophosphorus insecticides and concerning the acylating action of organophosphorus insecticides on the cholinesterase of insects as the chemical mechanism of this inhibition.

Insofar as at this time it is probable (1) that the object of phosphorylation in this ensyme is the hydroxyl group of serime which enters the polypeptide chain of its molecules, the working hypothesis was

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refined in that the selection was made among organophespherus compounds which are capable of phospherylating the alceholic hydroxyl group having some noticeable but weak acidic properties. Such organophosphorus compounds may be the anhydrides of acid esters of phosphorus acids, acid halides, substances similar to acid halides but containing, in the melecule, some pseudohaloid groups instead of the halogans, and, finally, neutral esters of acids of phosphorus in which one of the alkyl groups would correspond sufficiently well to an acidic alcohol. Since the acid halide and the anhydride derivatives of acids of phosphorus, as well as phosphorus esters of phenols, to which one may assign, without stretching the point, some functions of mixed anhydrides, have been already studied in detail and are being intensively studied in other laboratories, we concentrated our attention on the derivatives of acids of phosphorus which have residues of sufficiently acidic alcehols or mercaptans.

Further, we refined our working hypothesis in the sense that we realized the necessary presence in the molecule of an effective insecticide of a certain eptimum amount of the phosphorylating properties. Organophosphorus compounds with sharply displayed phosphorylating action as for instance dialkyl chlorophosphates, naturally cannot be sufficiently selective agents since they would show their phosphorylating action with any substrate of the erganism: unspecific proteins, carbohydrates and, figally, water. Such substances would be very rapidly detexified in the organism and would not reach those vitally important centers, the phosphorylation of which would be the consequency fellowing texic effect. On the other hand substances having but feeble phospherylating properties such as neutral esters of phosphoric or thiophosphoric acids, although able to reach the necessary receptors without large lesses, would be able to react then with the latter so slowly, if at all, that no action would result.

Therefore in the process of synthetic searches, when we found a poorly physiologically active substance, as an insecticide, with weak phosphorylating properties (slow hydrolysis, reactions with alcohols, amines), we along along the path of such a change in the structure of the molecule as to strengthen these properties. Conversely, if an insufficiently active (as an insecticide) substance turns out to be a good phosphorylating agent, we went along the reute of such a change of its structure which would yield an analog which would be less active.

The above shown principles were naturally not adhered to very firmly. The chemical distinction between the true cholinesterase, the imactivation of which assures the poisoning of warm blooded animals, and the

Approved For Release 2009/07/09: CIA-RDP80T00246A003900400002-3 p'seudo-cholinesterase or the insect cholinesterase is still not clear. We know too little about the mechanism of texic action to depend with complete confidence on the accepted working hypothesis. However, by being guided by these principles we built in general lines the plans for our syntheses and some positive results which we obtained were too systematic to be purely the work of chance.

Many literature data permit us to conclude that there is sense in searches in the area of thisphosphate compounds. The latter are distinguished from their oxygen analogs by lower texicity to warm blooded animals with preservation of the effective properties in respect to the insects.

In order to run the syntheses of substances with various structures it was necessary first of all to develop reasonable methods of proparation of the starting materials.

The majority of the known methods of synthesis of thie-erganephosphorus compounds are mainly based on the chloride derivatives of phosphorus. The latter are often inconvenient in handling and during the work with them it is necessary as a rule to conduct out of the sense of the reaction, by some means, the hydrogen chloride which forms in the reaction.

This-organophospherus compounds can be prepared also from the sulfides of phosphorus. Sulfides are sheap, simple in handling and their production is well in hand. Therefore we stopped with the sulfides of phosphorus as the initial starting base for the syntheses of this genophosphorus compounds.

Back in 1950 we explored the reactions of phosphorus sulfides  $P_4^8_5$ ,  $P_4^8_6$  and  $P_4^8_7$  with alcohols (2)

$$P_4 = RQH = (RO)_2 P(S)H + (RO)_2 P(S)SR$$
 $P_4 = RQH = (RO)_2 P(S)H + (RO)_2 P(S)SH + (RO)_2 P(S)SH$ 

It had been established that lower sulfides of phospherus P<sub>4</sub>S<sub>5</sub> and P<sub>4</sub>S<sub>5</sub> react with alcehols forming dialkyl thiophosphites and trialkyl dithiophosphates. The phospherus sulfide P<sub>4</sub>S<sub>7</sub> forms in its reaction a mixture of three substances which may be separated in a usual fractional distillation into dialkyl thiophosphites, dialkyl dithiophosphates and trialkyl dithiophosphates. The only exception is the reaction of this sulfide with methyl alcohol as the result of which there are formed but two compounds: dimethyl thiophosphite and trimethyl dithiophosphate. As to the mechanism of the reaction of the lower sulfides of phosphorus

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with alcohols, this appears to be a complex one. We examined the reaction of the sulfide  $P_4S_7$  with ethyl alcohol: along with the three products indicated above, there are also formed hydrogen sulfide, phosphine and phosphorus. It is very probable that all these products form as the result of not a single reaction but of several which run in parallel. The hydrolysis of  $P_4S_7$  is similarly emplex in its course.

We also studied (4) the previously known (5) products of the reaction of phosphorus decasulfide  $P_4 s_{10}$  with alcohols:

PAS10 + SROH + 4 (RO)2P(S)SH + 2H2S.

In this way there were developed the preparatively accessible methods of synthesis starting with phospherus sulfides of dialkyl thiophosphites, derivatives of trivalent phospherus which were obtained in this work for the first time, dialkyl dithiophosphates and trialkyl dithiophosphates. Here were prepared for the first time in pure state the free dialkyl dithiophosphates.

The constants of the empounds propared by us in those three classes of substances were used by us for the calculation of stemic refractions of sulfur bound to the phospherus in the thiene and the thiele type of linkage (6,7). The knowledge of the stemic refraction of sulfur was later of considerable help in the synthesis of compounds of product structures.

These three classes of compounds were the first objects tested for insecticidal action. All turned out to be weakly texts to insects. Their tests, as with all subsequent objects, were run on adult specimens of Eurygaster taken from the mite of hibernation, and on the post of citrus plants - the Marine flour were. In the system of memoralature used in the present report (table 1) one plus means that 95-100% of dead and paralyzed insects were observed after the action of the compound taken in concentration of 0.3% or greater. Two pluses correspond to the minimum lethal concentration of 0.1%, three pluses - 0.05%, etc. as shown in the table. The results of tests of the first three groups of compounds are shown in tables 2, 3 and 4.

As can be seen, dialkyl thisphosphites, dialkyl dithisphosphates and trialkyl dithisphosphates were poorly effective substances. From the dialkyl dithisphosphates we prepared and bested their petassium and nickel salts. These were also peer insecticides. If one compares these three classes of compounds, the most effective of them are dialkyl thisphosphites, derivatives of trivalent phosphorus.

Table 1.

Insecticid	al action o	r org	ganophosphorus	compounds	••	issects	(95-100%	kill).
			•	-4 -4 4				

encentr	16100,	>	. HOEAEIG
0.3	•	,	•
0.1			** .
0.05			+++
0.005			++++
0.0005	•		****

Table 2
Constants and insecticidal setion 66 tested preparations

No.	Prop	ration Formula	g.re. and	440	Action
1	M-1	(MoO),PMS	52-3/17 1.470	1.1892	. •
2	M-4	(EtO),PMS	67-8/12 1.459	7 1.0828	•
3	M-7	(Pr0), PMS	62-3/3 1.45	1.0390	. <b>-</b>
4	M-9	(100-Pr0) <sub>2</sub> PMS	. : 49-50/3 1.45	11 1.0135	•
5	M-13	(BuC),PES	89-90/4 1.45	B3 0.9974	•
6	M-15	(CH, CHCH, 0), PHS	77-8/6 1.49	1.0894	

The work on the synthesis of insecticides proper was begun with the prepartion of the simplest derivatives of dithiophespheric acid. There were prepared disulfides of the general formula (RO)<sub>2</sub>P(S)SSA, where A may be the simplest alkyl radical or a residue of dithiophespheric acid. The former were prepared by the reaction of salts of dialkyl dithiophespheres with sulfemyl chlorides (6) (this study was perfermed with the participation by the collaborator of our laboratory S.I.Godyny):

 $(RO)_{2}P(S)SM + R'SC1 + (RO)_{2}P(S)S-SR' + MC1$ 

2(RO)<sub>2</sub>P(S)SM + I<sub>2</sub> - (RO)<sub>2</sub>P(S)S-SP(S)(OR)<sub>2</sub> + 2 MI, while the latter compounds were prepared by exidation of the same salts by a solution of iodine (8). The resulting disulfides were tested both in the form of emulsions as well as dusts. However they also turned out to be rather weak insecticides (table 5). The compounds cited were but weak phosphorylating agents. They are very stable, are hydrolyzed with difficulty and are sufficiently stable in respect to alkalies. "e had to go on thus to the more labile anhydride forms.

As such we selected the acyl derivatives of dialkyl dithiophosphoric acids. These were prepared in good yields from sodium, petassium and lead salts of the corresponding acids and acyl halides (7).

- $(RO)_{2}P(S)SM + R^{1}COC1 + (RO)_{2}P(S)SCOR^{1} + MC1$

Table 3 \* Chastants and insecticidal action of the prepared substances

85,	frepn.	Formula	B.pt C p in mm	M.pt.	n <sub>D</sub> <sup>20</sup>	d <sub>4</sub> <sup>20</sup>	Action
	M-3	(MeO) <sub>2</sub> PSSH	62-3/5	-	1.5343	1.2888	-
2	M-5	(EtO) PSSH	81-2/5	-	1.5070	1.1654	. •
•	M-8	(PrO)2PSSH	85-6/3	-	1.4987	1.1040	•
4	M-11 (	iso-Pro) PSSH	71-2/3	-	1.4918	1.0911	+
)	M-36 (	iso-Bu0) 2PSSH	93/4	-	1.4889	1.0558	-
5	- (	EtO) PSSK	•	194-5	•	•	. +
7	- [(	Eto), PSS), ]Pb	•	75-6	•	•	-
c	- '[(1	Me0) <sub>2</sub> PSS] <sub>2</sub> N1	-	124-5	-	•	+

Table 4

Co	nstant.	and insecticidal	action of		ed compounds	
Nc.	Prepn.	Formula	B.pt C p mm.	n <sub>D</sub> <sup>20</sup>	d <sub>4</sub> <sup>20</sup>	Action
1	M-2	(MeO) <sub>2</sub> PS(SMe)	101/17	1.5292	1.2415	-
2	M-6	(BtO) PS(SEt)	115/10	1.5013	1.1168	+
4	M-9	(PrO) <sub>2</sub> PS(SPr)	115-6/3	1.4955	1.0561	-
4	M-12 (	100-Pr0) PS(SPr-1	so) 91-2/3	1.4843	1.0351	•
ζ		teo-Buch PS(SBn-4)			1 0150	4

Table 5

C	onstar	ts and insecticida	al action o	f tested	compour	nds.	
No	. Prep	on. Formula	B.pt.	M.pt.	n <sub>D</sub> 20	d420	Action
1	M-44	(EtO) <sub>2</sub> PS <sub>3</sub> Me	101-2/3	•	1.5500	1.2142	+
2	M-48	(Eto)2PS3Et		-	1.5431	1.1810	•
3	M-49	(EtO) <sub>2</sub> PS <sub>3</sub> Bu	139-40/4	•	1.5306	1.1246	+
4	M-43	(iso-Pro) <sub>2</sub> PS <sub>3</sub> Me	99-100/2	-	1.5297	1.1471	+
5	M- 50	(iso-Pr0)2PS3Et	117-8/3	-	1.5240	1.1189	+
6	M-52	(iso-Bu0)2PS3Et	129-30/3	-	1.5183	1.0866	+
7	M-16	$[(Me0)_2PSS]_2$	•	51-2	-	-	-
8	M-1#	[(EtO) <sub>2</sub> PSS] <sub>2</sub>	-	28-9	-	-	.=
		[(iso-Pr0) <sub>2</sub> PSS] <sub>2</sub>	-	91-2	-	-	-

It is of interest to note that the acetyl derivatives are obtained with equal success from the potassium and sodium, as well as lead salts of dialkyl dithiophosphoric acids. The benzoyl derivative may be prepared only from the lead salt. In turn the carbonate derivatives could be alkalimed only from the salts of the alkalimetals. As it is evident from table 6, the acyl derivatives prepared by us turned out to be more effective poisons, but still left much to be desired.

Even more active phosphorylating agents should be the oxygen analogs of the above cited acyl derivatives. These are constructed like the above compounds but contain one less atom of sulfur than the corresponding derivatives of dialkyl dithiophosphoric acids:

(RO)<sub>2</sub>P(S)OAcyl (RO)<sub>2</sub>P(O)SAcyl however, the chemistry of these compounds is but poorly explored. The synthesis of acyl derivatives from the salts of dialkyl thiophosphoric acids required special work on clarification of the reactivity of these. The results of this work may be expressed by the following scheme:

$$(RO)_{2}P(S)OM \rightarrow (RO)_{2}P(O)SR \\ \rightarrow (RO)_{2}P(S)OCOR' \\ \rightarrow (RO)_{2}P(O)SCOOR'$$

Table 6

C	onstants and effective	reness of action of		ivatives	
No.	Prepn. Formula	B.pt.	<b>n</b> D 20	d <sub>4</sub> <sup>20</sup>	Action
3	M-25 (Et0) 2PS 2COM	98/2	1.5154	1.1898	+
2	M-26 (180-Pr0) PS2C	000-1/3	1.4979	1.1177	-
3	M-,8 (180-Bu0) 2PS2CO	OMe 125-6/3	1.4929	1.0793	•
4	M-39 (iso-Pr0) PS2C	OPh m. 51-2		-	-
	M-27 (EtO) PS CO Me	105-6/4	1.5063	1.2171	<b>+</b>
(	M-29 (EtO) PS CO B	115/4	1.5001	1.1891	+
•	M-30 (Et0)2PS2CO2P1	125/6	1.4981	1.1620	+
S	M-34 (Et0)2P52C02B	162-3/6	1.4889	1.1281	-
•	M-31 (Et0) PS CO A	137-8/4	1.4925	1.1179	-
1.0	M-32 (iso-Pr0)2PS20	CO <sub>2</sub> Et 109/3	1.4894	1.1301	-
11	M-33 (iso-Pro)2PS20	CO <sub>2</sub> Am 138/3	1.4841	1.0773	•

During the study of the reactions of alkylation and acylation of the sodium, potassium and silver salts of dialkyl monothiophosphoric acid we established the dual character of reaction of these salts. In the reaction of alkyl halides (8) the alkylation occurred at the sulfur atom (see the reaction scheme), while in acetylation -at the oxygen atom (9). The

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By means of the infrared spectra we established the structure of the aikali and the silver salts of dialkyl monothiophosphoric acids. For the alkali salts we confirmed the data of Gere who had found that they have the thiono structure. The spectra of the silver salts were determined by us for the first time. These speak of the thiolo structure of the salt with strong association of the molecules]

The data concerning the structure of the salts in the absence of their tautomerism which is apparently improbable in this case speak of the fact that their reactions of alkylation and acylation may proceed with or without the transfer of the reactive center. This study was made in connection with work in the area of tautomerism and reactivity of organophosphorus compounds and is described in more detail in the report by M.I.Kabachnik.

Tests of the acyl derivatives of thiolophosphoric acid showed that there are weak insecticides. Similarly poorly effective turned out to be some of the alkyl derivatives (table 7). Strong insecticidal properties are had in the carbonate derivatives[ their effectiveness is such that they cause a 95-100% kill of the flour worm after use in the concentration 0.05% or greater (table 7, substances 6-7).

We also prepared mixed alkyl phosphoric disulfides from the corresponding sodium salts of dialkyl monothiophosphoric acid

 $(RO)_2P(S)ONa + R'SC1 \rightarrow (RO)_2P(O)S-SR' + MaCl.$ Since sulferly chlorides (8). The thus obtained derivatives showed high toxicity to the Eurygaster insect, especially substance M-51. For us this was a certain degree of success, although substance M-51 is less active than Tiofos in its contact action.

Thus, it seemed that monothiophosphoric derivatives give more chances for success in finding practically interesting substances. However, we did not consider it rational to develop this area insofar as it is being studied in many other laboratories.

A further study was directed along two main paths: 1) preparation of derivatives of the simplest alkyl-thiophosphonic acids and 2) the synthesis of trialkyl dithiophosphates containing definite substituents in predominantly the  $\beta$ -positions.

Alkyl-thiophosphonic derivatives may, judging from literature data, be more powerful inhibitors of cholinesterase than the corresponding

thiophosphoric derivatives.

Among the derivatives of alkyl-thiophosphonic acids we studied the dalkyl esters of alkylthiolophosphonic and alkylthionophosphonic acids (13). These substances were prepared by the action of alkyl halides on dialkyl thiophosphite sodium salt. For preparation of the thiolo derivatives, the thiono forms were isomerized by heating in a sealed tube in the presence of alkyl halides, according to Pishchimuka method. The thiolo derivatives were also prepared from the appropriate halides of alkylphosphonic acid and alsohol or mercaptan, as well as by alkylation of the alkali salts of esters of alkylthionophosphonic acids (this work was carried out with participation of the degree aspirant N.I.Kurochkin and E.E.Kugucheva).

$$P_4S_6 + ROH + (RO)_2P(S)H$$
 $R^*PSC1_2 + ROH$ 
 $R^*P(S)(OR)_2$ 
 $R^*P(S)(OR)_2$ 
 $R^*P(S)(OR)OH$ 
 $R^*POC1_2$ 
 $R^*P(O)(SR)OR$ 
 $R^*P(S)(OR)ONa$ 
 $R^*P(S)(OR)ONa$ 
 $R^*P(S)(OR)ONa$ 

· Table #

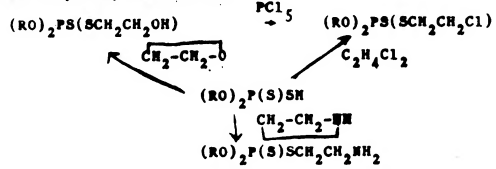
CO	necance	and insecticidat	action of the	Detes		
No.	Prepn.	Formula	B.pt.	<b>n</b> 20	d <sub>4</sub> <sup>20</sup>	Action
1	K-2	HePS(OBt)2	76-8/13	1.4610	1.0553	•
2	K-19 .	EtPS(OMe)(OBt)	74-5/10	1.4665	1.0647	-
3	M-21	EtPS(OEt) <sub>2</sub>	80-3/10	1.4576	1.0324	-
4	K-3	PrPS(OEt)2	64-6/2	1.4596	1.0158	-
5	K-4	BuPS(OBt)2	74-7/2	1.4600	1.0004	-
6	K-5	PhCH2PS(OBt)2	123-6/2	1.5305	1.1031	<b>+</b>
7	K-10	MePS(OBu)2	72-4/1	1.4535	0.9843	-
3	K-11	EtPS(OBu)2	80-3/2	1.4533	0.9775	•
1	K-22	FtPS(OEt)(OCOMe)	68 - 9/1.5	1.4701	1.1232	+

The first representatives of this group of substances were prepared by as back in 1950(M-24). The thiono and the thiolo esters (tables 8-9) differ considerably in the insecticidal properties. While the thiono esters, as should be expected, were only rather weak insecticides, the thiolo derivatives turned out to be quite effective. Thus, substances M-24, K-6 and K-9 were quite close to Tiofos.

By this path we first reached effective insecticides which along with the contact action also possessed the systemic action. However, none of the compounds possessed properties superior to the known insecticides in use now.

Somewhat different considerations lay in the foundation of the second path. I remind you that disulfide derivatives turned out to be too stable while the acyl derivatives, conversely, were too labile in order to make good insecticides. We stopped on the trialkyl dithiophosphates which carry substituents in the alkyl group connected with sulfur, especially in the  $\beta$ -position. We expected of these derivatives the optimum phosphorylating ability. Naturally only the experimental method could be used to prove this expectation.

Thus, by the reaction of ethylene oxide with dialkyl dithiophosphates we ( together with the degree aspirant at Y.N.I.I.V V.N.Odnoralova) prepared β-hydroxy derivatives



and from these we prepared  $\beta$ -acetoxy and  $\beta$ -chloro derivatives (11). We also prepared  $\beta$ -aminoethyl dithiophosphate, previously prepared in V.I.Z.R. in the A.I.Kulikev's laboratory. There were prepared other  $\beta$ -amino esters, as well as their derivatives. If one compares the activity effectiveness of compounds shown in table 10 it is easy to see that the introduction of substituents in the  $\beta$ -position increases the insecticidal properties in this series apparently along with the electronegativity of the substituents. The most active was the chloro derivative which had been prepared by Schrader (12).

The next step should be the synthesis of sulfur analogs

(RO)<sub>2</sub>P(S)SCH<sub>2</sub>CH<sub>2</sub>SR'
especially since at that time there had appeared the patent of Schrader
(13) on Systox, which is a β-substituted derivative of monothiophosphoric
acid. At first it was proposed to use for the synthesis of β-alkylmercapto substituted compounds the reaction of addition of dialkyl dithiophosphates to vinyl thio ethers (this work was carried out in collaboration with Coworkers of Institute of Organic Chemistry M.F. Shostakovskii
E.N.Ir:lezhaeva and N.N.Uvarova).

Table 9

	Constan	ts and insecticidal	action of			
No.	Prepn.	Formula .	B.pt.	<b>n</b> <sub>D</sub> <sup>20</sup>	<b>d</b> <sup>20</sup> <sub>4</sub>	Action
1	K - 9	MePO(SEt)(OBt)	106-8/18	1.4718	1.0904	+++
2	Ku-1	MePO(SBt)(OPr)	71-2/3	1.4510	1.0356	+
3	Ku-3	MePO(SEt)(OPr-iso)	60-2/1	1.4749	1.0555	++
4	Ku-2	MePO(SEt)(0Bu-1so)	69-70/1	1.4666	1.0222	++
5	K-23	MePO(SCH <sub>2</sub> OMe)(OBt)	117-20/2	1.4778	1.1538	. +(+)
¢	K-18	EtPO(SMe)(OBt)	93/9	1.4790	1.1058	+(+)
7	MDS-24	EtPO(SEt)(OEt)	66-8/2.5	1.4747	1.0670	+++
8	K-12	EtPO(SBu)(OBu)	92-4/2	1.4660	0.9951	•
9	K-6	PrPO(SEt)(OEt)	85-6/3	1.4733	1.0447	++
10	K-7	BuPO(OEt)(SEt)	98-100/3	1.4728	1.0262	+
11	K-8	PhCH2PO(SEt)(OEt)	134-6/2	1.5350	1.1263	+

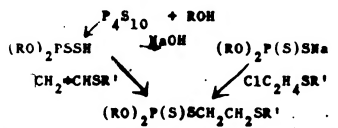
Table 10

	Consta	ints and insecticion	dal action (	of the com	pounds test	be
No.	Prepr	. Formula	B.pt. at 2 mm.	n <sub>D</sub> <sup>20</sup>	d <sub>4</sub> <sup>20</sup>	Action
1	MO-5	(MeO) 2PS2C2H4OH	-	1.5380	1.2911	+
2	MO-1	(Eto) PS2C2H4OH	119-20	1.5250	1.2042	<b>+</b>
3	MO-2	(Pr0),PS,C,H,OH	124-6	1.5140	1.1440	+
4	MU-4	(iso-Pr0)2PS2C2H4	он 118-9	1.5083	1.1323	•
5		(iso-Bu0)2PS2C2H4		1.5045	1.0965	+
0		(EtO) PS2C2H4OAc	135-6	1.5010	1.1845	++
7		(iso-Bu0) PS2C2H4	OAc 140-1/1	1.4890	1.0948	+
8		(Et0),PS,C,H,C1	103-4	1.5230	1.2270	+++
9	MO-14	(EtO) PS_C_HANH	-	1.5287	. •	•
10	MO-19	(Pro),PS,C,H,NH,	-	1.5130	1.1099	-
11	MO-21	(1so-Bu0), PS, C, H,		1.5065	1.0705	-
1 2	MO-18	(EtO) <sub>2</sub> PS <sub>2</sub> C <sub>2</sub> H <sub>4</sub> MH <sub>2</sub> (picrate)	•	-		-

However it appeared that under the usual conditions there are not formed maximum the  $\beta$ - but the  $\alpha$ -derivatives, the structure of the addition products being determined by a known synthesis from  $\alpha$ -chlorodiethyl sulfide as shown in the following scheme, and by other means (14).

The substances prepared were tested on the Eurygaster insects. They turned out to be weak insecticides against this insect (table 11).

The synthesis of the first representatives of β-alkylmercapto substituted trialkyl dithiophosphates was achieved by us in 1953 in good yield of about 90-5% starting with sodium or potassium salts of dialkyl dithiophosphoric acids and β-chloroalkyl sulfides:



The latter were prepared in the beginning of the study by the reaction of hydrogen chloride with products of addition of mercaptans and vinyl ethers. The substances proved to be highly effective insecticides with contact-systemic action, exceeding by many orders the eisting organophosphorus chemical poisons (table 12). Therefore in the course of the following study we found a new and better method of synthesis, based on ethylene oxide, ethyl mercaptan and dialkyl dithiophosphoric acid.

Table 11

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Co	nstant	s and insecticidal p	roperties of	•		ted
No.	Prepr	. Formula	B.pt.	. nD	$\mathbf{d}_{A}^{20}$	Action
1	M-65	(EtO) PS2CHMeSEt	109-10/3	1.5290	1.1392	+
2	M-54	(BtO) PS2CHMeSBu	109-10/2	1.5198	1.0965	+
3	M-59	(BtO) PS CHMoSC NA	Bu 123-5/3	1.5125	1.0940	+
4	M-56.	(1sombu0) PS, CHOto SE	113-5/2	1.5070	1.0556	+
5	M-55	(1so-Bu0) PS_CHMeSBu	121-2/2	1.5070	1.0384	<b>+</b>
6	M-53	(1so-Bu0) PS_CHOLOSC	H <sub>4</sub> 0Bu 124-6/3	1.5012	1.0422	+
7	M-86	(EtO) PS CH SPr	145-6/4	1.5270	1.1308	-
8	M-87	(BtO) PS_CH_SPr-1sc	133-4/4	1.5210	1.1312	•
ı	M-82	Table 12. (Me0) <sub>2</sub> PS <sub>2</sub> C <sub>2</sub> H <sub>4</sub> SMe	71-1.5/0.00	04 1.5580	1.2493	++++
2	M-81	(NeO)2PS2C2H4SEt	91-2/0.003	1.5598	1.2065	++++
3	M-80	(BtO) PS C H SNo	127-8/2	1.5405	1.1699	++++
4	M-74	(BtO) PS C H SEt	129-30/2	1.5350	1.1445	+++++
5	M-85	(BtO) PS2C2H4SPr	143-4/2	1.5275	1.1260	+++
6	M-77	(BtO) PS2C2H4SBu	150/1	1.5255	1.1040	++
7	M-75	(1so-Pro) 2PS2C2H4SEt	: 134-5/3	1.5189	1.0887	++
8	M-76	(iso-Pro)2PS2C2H4SB	148-9/4	1.5206	1.0622	+
	M-58	(Bt0)2PS2C2H4SC2H401	Su 188/3	1.5160	1.1050	+++
10	<b>M</b> ≠78	(1so-Pr0) 2P82C2H48C		1.5090	1.0967	+

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E.N.Prilezhaeva also realised the addition of dialkyl dithiophosphates to thiovinyl ethers contrary to the Markovnikov rule in the presence of isopropylbensene hydroperoside. Here there were formed \$-substituted trialkyl dithiophosphates in 50-60% yields.

Tests of properties of compounds of this type maxima were conducted mainly with the specimen M-74. These tests were run on various plant cultures with several species of insects. Substance M-74 turned out to be a highly effective poison against the red apple mite, and at concentration of 0.05% protected the apple trees for up to a month and a half. M-74 remained effective up to two months against the Tetranychus mite on roses. At this concentration substance M-74 was effective up to three weeks against the green plum aphid. The duration of action of the substance in many experiments was greater than for Merkaptofos on the same objects.

Especially promising is the preceding treatment of summer wheat grains. Seeds treated with 2% solution of the M-74 concentrate ( with 30% content of the active substance) yield sprouts which are poisonous to Burygaster for 2-3 weeks (100% kill). The norm of consumption of the substance is 120-150 grams of the active principle of M-74 per hectare of the field which fact shows the highly economic feature of M-74\*

Interesting data, requiring further tests, were obtained in control of pests of sugar beet, corn, etc.

Not less interesting are substances M-81 and M-82. Their study was run amaller scale than that of M-74. These are also contact-systemic insecticides. In effectiveness they are very close to M-74. However these empounds are weakly toxic to warm-blooded animals in comparison with aptofos or M-74. On intravenous application to rabbits they are almost times less toxic than Tiofos. Thus, as the result of the study we d in finding a group of insecticides - \$-alkylmercaptoethyl esters 'kyl dithiophospheric acid, which have high insecticidal and activity with contact and systemic action.

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## Remarks

to (B.E.Arbuzov Chem.Inst., Kazan Section, Academy of for

The Mastryukova illustrated well the meaning of synthesis apphorus insecticides on the basis of phosphorus sulfider state are is very cheap and in addition has great possibilities for diverse synthesis.

sis made in the area of thiophosphoric compounds are only in these studies.

Yakovlev (Brain Institute, Acad. Med. Sciences USSR)

ion of active centers of enzymes. At this time the process in this area. The results given by T.A.Mastryukova justification of development of the previous study of the action of extrasic substances. For the reaction of insecticides with the phosphorylation reaction must proceed. For clarification has to which functional groups in the molecule of englishing phosphorylated, modern means of study were used (chromatic phosphorylated, modern means of which it was possible to the phosphoric acid in yield of 40%. This is a proof that in of cholinesterase the hydroxyl group is phosphorylated. If some as such, its hydroxyl group does not react with organic asecticides. It is necessary to say that all these studies evelopment stage.

Mazar and Bodansky it is known that the mechanism of pure state of true and pseudo cholinesterases differs. Therefore the cholinesterases. In addition it is desirable to show in the table of ratio of action of the insecticide and in the table of ratio of action of the compounds on warm blooded animals and on a letter ratio would be of more use than insecticidal action alone.

N.T.Mel'nikov (Pertilizer and Insect of ungicide Res Jost and

N.I.Mel'nikov (Pertilizer and Insectofungicide Res. Inst. nem. after Ya.V. Samoilov, Moscow)

I have no confidence in the fact that biologists consider that seaticide acts only on chelinesterase. It seems to me that the cannot be solved in such single-valued manner. There are substituted inhibit cholinesterase and at the same time are not insectioned example such is the series of quaternary ammonium bases. There is of synerals raised by A.E.Arbusov: addition of small amounts of substances semetimes aid the elevation of toxicity by 2-3 fold. The true of DDT and other insecticides. At this time the study of same tures is being carried out very extensively.

V.A.Yakovlev (Brain Institute, Academy of Med. Sciences U:
The problem of antagonism and synergism of organophosphorus insecticides is not being worked on sufficiently. One should request the BiologInstitute of the Academy of Sciences USSR to occupy itself with this pro-